NEW DRUG MANUFACTURING LICENSE APPLICATION PLEASE COMPLETE THIS FORM FULLY—INCOMPLETE APPLICATIONS WILL BE RETURNED See Page 2 for Instructions.

☐ NEW APPLICANT [RELOCATION	OWNERSHIP CH	HANGE 🗌 OWNE	RSHIP AND LOCATIO	N CHANGE	
1. Name of Firm			Facility Operator (name ar	nd title)		
DBA (List additional DBAs on separate sheet if necessary.) 10			Facility Telephone Numbe (er 11. Faci	lity FAX Number	
3. Facility Address (number, street))	12.	24-Hour Emergency Telep	phone Number 13. E-M	ail Address	
4. Facility Address (continued)		14.	Correspondent (name and	d title)		
5. City	State 2	ZIP Code 15.	Correspondent Telephone	Number 16. Corr	espondent FAX Number	
6. Mailing Address (if different or P	.O. Box number)	17.	Country (if other than United	ed States) 18. FDA	CFN or FEI Number	
7. Mailing Address (continued)			19. Web site (URL)			
8. City	State Z	ZIP Code 20.	Interstate Commerce Product Shipped	☐ Product or Raw I	Materials Received \ N/A	
 Type of Ownership ☐ Individual/Sole Proprietor 	ship	·	nited Liability Company	☐ Nonprofit ☐	Other:	
22. Corporate Name (if applicable)		Star	State of Incorporation			
23. Owners' or Officers' Names and	Titles	Owi	Owners' or Officers' Names and Titles			
24. Size of Facility (square feet):			mber of Employees at th	nis Facility:		
 Stage of Manufacture at Date of Manufacturing products 	Application (check all that ap	• • •	Plant construction/design (Targeted Completion Date:)			
☐ Validation – Completion I		Oth	er (specify):			
26. Intended Drug Destination (chec						
Commercial distribution 27. Type of Drug Product (check all	Human clinical trial	ls (investigational use)	oution only U.S.	distribution	
☐ Prescription* ☐ Over-	-the-counter		ion or Both is checked	d refer to PDMA requir	ements on instruction page 2.	
28. Drug Products Manufactured at 700 Bulk pharmaceutical 701 Medical gases 702 Radioactive 703 Veterinary	s (API)	d substances DEA #: _)	07 Biotech 08 Biologics 09 Parenteral 10 Oral Dose (solid/liqui	☐ 711 Pre-IND ☐ 712 Topical ☐ Other (specify): d)	
-	tivities employed or planr	ned in the manufactur	e of the drugs listed abo	ove. Indicate if these p	rocesses/activities will be done at	
Processes/Activiti Aerosolization Aseptic		Contract	· · · · · · · · · · · · · · · · · · ·	ocesses/Activities	In-house Contract	
Coating Emulsification Encapsulation Fermentation/tissue culture	□ □ □ viral □		Repackage Only Sterilization Suspension Tableting			
vector/gene therapy Liquid Mixing			Other (Specify):			
30. Payment Codes (Check only A—\$1600 B-	/ one code—see page 2 f -\$850	or schedule)	31. License Fees Due a. License Fee (se		Enter Each Fee Below:	
MAKE CHECKS PAYABLE TO: DEPARTMENT OF HEALTH SERVICES See page 2 for mailing address				f Applicable – see page	· · · · · · · · · · · · · · · · · · · 	
		c. Total Payment		th and Safoty Codo, \$111630		
The Food and Drug Branch MUST BE NOTIFIED of any change in the application information as provided by CA Health and Safety Code, §111630. By signature, I declare under penalty of perjury that all information provided herein is true and correct.						
32. Signature	Printed Name	inomiadon provid	Title		Date	
			ITE BELOW THIS LINE			
License Number	Expiration Date	Date Received	Payme	ent Type	Amount \$	

NEW DRUG MANUFACTURING LICENSE APPLICATION INSTRUCTIONS

A separate application is required for each place of business. Please complete and/or amend this application as is most appropriate to your facility. Include the appropriate fee for each application and make payable to: <u>DEPARTMENT OF HEALTH SERVICES</u>. The fee must accompany this application or it cannot be processed. Unsigned or incomplete applications cannot be processed. The following are further instructions on how to complete this application:

New Applicant: Place an (X) in the box next to New Applicant if your firm has not previously applied for a Drug Manufacturing License at this location while under the current ownership. **This license is non-transferable.** If your firm has changed location, ownership, or both, place an (X) in the appropriate box **and also** in the box next to New Applicant. For any section that does not apply to your company, please indicate with (N/A). Do not leave any sections blank.

- 1. Name of Firm: Enter full name of business, corporation, company, or organization applying for licensure.
- 2. **DBA:** Enter any other name(s) your company is doing business as.
- 3.-5. Facility Address: Enter the number, street, city, state, and ZIP code for this facility location.
- 6.-8. Mailing Address: Enter the full mailing address if different from the facility address.
 - 9. Facility Operator: Enter the full name(s) of the person(s) in charge of drug manufacturing at this facility and their title(s).
 - 10. Facility Telephone Number: Enter daytime business telephone number of this facility.
 - 11. Facility FAX Number: Enter facility FAX number.
 - 12. 24 Hour Emergency Telephone Number: Enter telephone number to be called in the event of an emergency.
 - 13. E-mail Address: Enter facility e-mail address.
 - 14. Correspondent: Enter the name of the person to contact for information regarding this application and their title.
 - 15. Correspondent Telephone Number: Enter the daytime business telephone number of the contact person.
 - 16. Correspondent FAX Number: Enter the daytime business FAX number of the contact person.
 - 17. Country: Enter the country where your facility is located, if outside of the United States.
 - 18. FDA CFN or FEI: Enter your US Food and Drug Administration Central File Number or Federal Establishment ID, if known.
 - 19. Web site: Enter the Web site address for your business, if applicable.
 - 20. **Interstate Commerce:** Place an (X) in all appropriate boxes that correctly describe your business' receipt or distribution of products or materials through or into interstate commerce.
 - 21. **Type of Ownership:** Place an (X) in the box next to the appropriate legal description of the facility's ownership.
 - 22. Corporate Name: Enter corporate name if applicable. Enter state of incorporation if applicable.
 - 23. Owners' or Officers' Names: List the business owners' or officers' names and titles. USE ADDITIONAL SHEETS IF NECESSARY.
 - 24. Size of Facility: Indicate the most appropriate size (in square feet) at this facility and the approximate number of employees at the facility.
 - 25. **Stage of Manufacture:** Place an (X) in the box next to the stage of manufacture your products are in at the time of application submission. Check all that apply.
 - 26. Intended Drug Destination: Place an (X) in the box adjacent to the destination(s) for your manufactured products. Check all that apply.
 - 27. **Types of Products:** Place an (X) in each box that applies to the type of drugs manufactured or to be manufactured. For human prescription (Rx) drug manufacturers, refer to PDMA requirements below*.
 - 28. **Products Manufactured:** Place an (X) in the box adjacent to each product area box that applies to the drugs manufactured or to be manufactured. Use additional sheets if necessary.
 - 29. **Manufacturing Processes:** Place an (X) in the columns adjacent to all applicable processes to be performed in-house and/or contracted-out. Leave line blank if the indicated process will not be applied to the manufacturing of listed drugs. List additional processes or methods as needed herein or on additional sheets if necessary.
 - 30. Payment Fee Code: Your license fee is based on the application type, number of employees, amount of sales, and the type of drugs being manufactured at the facility.

Application Type	Fee	Payment Interval	Payment Code
New, Relocation, or Ownership Change	\$1600	First License only	А
New (**Special/Small Firms)	\$850	First License only	В

^{**} Special or Small Firm types are limited to companies that 1) repack medical gas only, OR 2) employ three or fewer people and have an annual sales of less than \$500,000.

- * PDMA (Prescription Drug Marketing Act) Requirements: If your firm manufactures human prescription (Rx) drugs, an additional \$100.00 must be added to the license fee and a Disclosure Statement (Form EH 53) must be submitted for each person listed on lines #9 and #23 (instructions provided therein). Information relevant to the PDMA, (e.g., Disclosure Statements and Applicant Fingerprint Live Scan requirements) can be reviewed at: http://www.dhs.ca.gov/fdb/HTML/Drug/PDMA.htm.
- 31. License Fee Due: Enter appropriate fees due.
 - a. Enter license fee according to payment codes in #30.
 - b. Add \$100 PDMA fee if it applies to your firm. See PDMA requirements above*.
 - c. Enter Total Payment Due by adding a and b.
- Sign the application, print your name, print your title, and enter the date. All signatures must be original.

Make checks payable to: <u>DEPARTMENT OF HEALTH SERVICES</u> Mail Application and Check to: (below)

Regular Mail: California Department of Health Services

Food and Drug Branch - Cashier

MS 7602

P.O. Box 997435

Sacramento, CA 95899-7435

Overnight Mail: California Department of Health Services

Food and Drug Branch - Cashier 1500 Capitol Avenue, MS-7602

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Sacramento, CA 95814

If further questions exist, please contact the FDB License Desk for Drug Manufacturing at (916) 650-6500, or visit our web site at: http://www.dhs.ca.gov/fdb/.

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